=> d his

(FILE 'HOME' ENTERED AT 22:05:35 ON 16 AUG 2006) FILE 'REGISTRY' ENTERED AT 22:05:41 ON 16 AUG 2006 L1 STRUCTURE UPLOADED L2 0 S L1 STRUCTURE UPLOADED L3 L40 S L3 L50 S L3 L6 0 S L3 FULL STRUCTURE UPLOADED L7 L8 11 S L7 801 S L7 FULL L9 FILE 'HCAPLUS' ENTERED AT 22:13:42 ON 16 AUG 2006 L10 4 S L9 2 S L10 AND GIBLIN, G?/AU L11 2 S L10 NOT L11 L12 0 S L12 AND HALL, A?/AU L13 0 S L12 AND HURST, D?/AU L14L15 0 S L12 AND KILFORD, I?/AN 0 S L12 AND KILFORD, I?/AU L16 0 S L12 AND LEWELL, X?/AU L17 0 S L12 AND TAYLOR, A?/AU L18 0 S L12 AND NOVELLI, R?/AU L19 FILE 'CAOLD' ENTERED AT 22:15:54 ON 16 AUG 2006

0 S L9

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L20

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                 The first reclassification of IPC codes now complete in
         JUN 02
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        JUN 26
                 and display fields
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=> file reg

10508761

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TOTAL SINCE FILE SESSION ENTRY 0.21

0 ANSWERS

0.21

FULL ESTIMATED COST

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STRUCTURE FILE UPDATES: 15 AUG 2006 HIGHEST RN 901654-60-2 DICTIONARY FILE UPDATES: 15 AUG 2006 HIGHEST RN 901654-60-2

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STRUCTURE UPLOADED Ll

SAMPLE SEARCH INITIATED 22:06:18 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -6753 TO ITERATE

2000 ITERATIONS 29.6% PROCESSED INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

ONLINE **COMPLETE** FULL FILE PROJECTIONS:

COMPLETE BATCH

PROJECTED ITERATIONS: 130134 TO 139986 0 TO PROJECTED ANSWERS:

L20 SEA SSS SAM L1

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L3STRUCTURE UPLOADED

=> d 13

L3 HAS NO ANSWERS

L3 STR

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Updated Search

Structure attributes must be viewed using STN Express query preparation.

=> s 13

SAMPLE SEARCH INITIATED 22:07:15 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 6753 TO ITERATE

29.6% PROCESSED 2000 ITERATIONS 0 ANSWERS

0 ANSWERS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

> BATCH **COMPLETE**

> > 139986

PROJECTED ITERATIONS: 130134 TO

PROJECTED ANSWERS: 0 TO

0 SEA SSS SAM L3

=> s 13

SAMPLE SEARCH INITIATED 22:07:20 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 6753 TO ITERATE

29.6% PROCESSED 2000 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 130134 TO 139986 0 TO

PROJECTED ANSWERS:

 L_5 0 SEA SSS SAM L3

=> s 13 full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 166.50 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y) /N or END:y FULL SEARCH INITIATED 22:07:27 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 134478 TO ITERATE

100.0% PROCESSED 134478 ITERATIONS 0 ANSWERS

SEARCH TIME: 00.00.01

L6 0 SEA SSS FUL L3

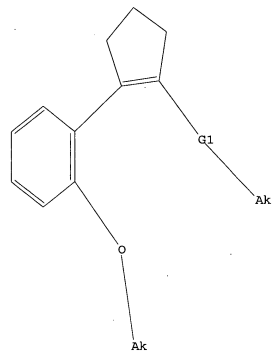
Uploading C:\Documents and Settings\brobinson1\My Documents\stnweb\Queries\34y.str

STRUCTURE UPLOADED L7

=> d 17

L7 HAS NO ANSWERS

L7 STR



G1 Ph,Hy

Structure attributes must be viewed using STN Express query preparation.

=> s 17

SAMPLE SEARCH INITIATED 22:13:34 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 6762 TO ITERATE

29.6% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS:

NS: 130310 TO 140170

PROJECTED ANSWERS:

378 TO 1108

L8 11 SEA SSS SAM L7

=> s 17 full

THE ESTIMATED SEARCH COST FOR FILE 'REGISTRY' IS 166.50 U.S. DOLLARS DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N or END:Y FULL SEARCH INITIATED 22:13:38 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 134677 TO ITERATE

100.0% PROCESSED 134677 ITERATIONS

801 ANSWERS

11 ANSWERS

SEARCH TIME: 00.00.01

L9 801 SEA SSS FUL L7

=> file hcaplus

Updated Search

COST IN U.S. DOLLARS

SINCE FILE TOTAL SESSION 338.72 338.93

FULL ESTIMATED COST

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s 19

L10 4 L9

=> s 110 and giblin, g?/au 58 GIBLIN, G?/AU

L11 2 L10 AND GIBLIN, G?/AU

=> d l11, ibib abs fhitstr, 1-2

L11 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:371223 HCAPLUS

DOCUMENT NUMBER:

142:430146

TITLE:

Preparation of cyclopentene compounds which bind with

high affinity to the EP1 receptor

INVENTOR(S):

Giblin, Gerard Martin Paul; Hall, Adrian;

Hurst, David Nigel; Kilford, Ian Reginald; Lewell,

Xiao Qing; Naylor, Alan; Novelli, Riccardo

PATENT ASSIGNEE(S):

Glaxo Group Limited, UK

SOURCE:

PCT Int. Appl., 201 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2005037793	A1	20050428	WO 2004-EP11364	20041006
W: AE, AG,	AL, AM, AT	, AU, AZ,	BA, BB, BG, BR, BW, BY,	BZ, CA, CH,
CN, CO,	CR, CU, CZ	, DE, DK,	DM, DZ, EC, EE, EG, ES,	FI, GB, GD,
GE, GH,	GM, HR, HU	, ID, IL,	IN, IS, JP, KE, KG, KP,	KR, KZ, LC,

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LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
             SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN, TD, TG
                                            EP 2004-765925
                                20060621
                                                                    20041006
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK, HR
PRIORITY APPLN. INFO.:
                                            GB 2003-23581
                                                                 Α
                                                                   20031008
                                            WO 2004-EP11364
                                                                    20041006
                         MARPAT 142:430146
OTHER SOURCE(S):
GI
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$$R^{8}$$
 R^{9} $F_{3}C$ N $CO_{2}H$ R^{2} R^{2} R^{3} I C_{1} C_{1} II

The title compds. I [A = (un)] substituted aryl, 5-6 membered heterocyclyl, AB bicyclic heterocyclyl; B = Ph, pyridyl; Z = O, S, SO, SO2; R1 = CO2H, CN, CH2CO2H, etc.; R21, R22 = H, halo, alkyl, etc.; R3 = (un)substituted alkyl (wherein 1 or 2 of the non-terminal carbon atoms are optionally substituted by (un)substituted NH, O, S, SO, SO2), alkenyl, etc.; R8, R9 = H, Cl, F, CF3, alkoxy, alkyl], useful in the treatment of conditions mediated by the action of PGE2 at EP1 receptors, were prepared Thus, hydrolysis of (2,4-dichlorophenyl) methyl 6-{2-[2-{[(2,4dichlorophenyl)methyl]oxy}-5-(trifluoromethyl)phenyl]-1-cyclopenten-1-yl}-2-pyridinecarboxylate with 2M NaOH solution afforded II. The compds. I had an antagonist pIC50 value of 6.0 to 9.5 at EP1 receptors. pharmaceutical composition comprising the compound I is disclosed. IT 850861-20-0P RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (preparation of 1,2-disubstituted cyclopentenes which bind with high affinity to the EP1 receptor) 850861-20-0 HCAPLUS RN

850861-20-0 HCAPLUS
2-Pyridinecarboxylic acid, 6-[2-[5-bromo-2-[(2,4-difluorophenyl)methoxy]phenyl]-1-cyclopenten-1-yl]- (9CI) (CA INDEX NAME)

CN

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 2 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:818387 HCAPLUS

DOCUMENT NUMBER:

139:323536

TITLE:

Preparation of [2-(2-alkoxyphenyl)cyclopent-1-enyl] substituted (hetero) aromatic carboxylic acids with

high affinity to the EP1 receptor

INVENTOR(S):

Giblin, Gerard Martin Paul; Hall, Adrian;

Hurst, David Nigel; Kilford, Ian Reginald; Lewell,

Xiao Qing; Waylor, Alan; Novelli, Riccardo

PATENT ASSIGNEE(S):

SOURCE:

Glaxo Group Limited, UK PCT Int. Appl., 180 pp.

CODEN: PIXXD2

10 508761

DOCUMENT TYPE:

Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	PATENT NO.		KIND		DATE \		APPLICATION NO.				DATE						
WO	2003	0849	17		A1		2003	1016	\	WO 2	003-1	EP36	61		2	0030	407
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		CO,	CR,	CU,	CZ,	DΕ,	DK,	DM,	ďΖ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	ΙL,	IN,	IS,	JŊ,	KE,	KG,	ΚP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK\	MN,	MW,	MX,	ΜZ,	NO,	NZ,	OM,	PH,
							SD,										
		UA,	UG,	(US)	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw						
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		BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	Gb,	GW,	ML,	MR,	NE,	SN,	TD,	TG
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	1492																
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PRIORIT			1							GB 2	002-	8Ò45		1	A 2	0020	408
			- 1								003-					0030	
			- 1							WO 2	003-1	EP36	61	1	N 2	0030	407
			- 1														

Updated Search

OTHER SOURCE(S):

MARPAT 139:323536

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The title compds. [I; A = (un) substituted Ph, 5-6 membered heterocyclyl, bicyclic heterocyclyl; R1 = CO2R4, CONR5R6, CH2CO2R4, alkyl, etc.; R2 = halo, alkyl, CN, etc.; R3 = alkyl wherein 1 or 2 of the non-terminal carbon atoms may optionally be replaced by NR4, O, SOn (n = 0-2), etc.; R4, R5 = H, alkyl; R6 = H, alkyl, SO2aryl, etc.; R8, R9 = H, alkyl; n = 0-2], useful for treating condition which is mediated by the action of PGE2 at EP1 receptors, were prepared E.g., a multi-step synthesis of [2-(5-chloro-2-benzyloxyphenyl)cyclopent-1-enyl]benzoic acid (starting from 1,2-dibromocyclopentene and (3-ethoxycarbonylphenyl)boronic acid), was given. The compds. I had an antagonist pIC50 value of between 7.0 and 9.5 at EP1 receptors and pIC50 value of <6.0 at EP3 receptors. Pharmaceutical composition comprising the compound I is claimed.

RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses)

(preparation of [2-(2-alkoxyphenyl)cyclopent-1-enyl] substituted (hetero)aromatic carboxylic acids with high affinity to the EP1 receptor)

RN 612832-50-5 HCAPLUS

3-Pyridinecarboxylic acid, 5-[2-[2-(phenylmethoxy)-5-(trifluoromethyl)phenyl]-1-cyclopenten-1-yl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d his

CN

(FILE 'HOME' ENTERED AT 22:05:35 ON 16 AUG 2006)

2

FILE 'REGISTRY' ENTERED AT 22:05:41 ON 16 AUG 2006

L1 STRUCTURE UPLOADED

L2 0 S L1

STRUCTURE UPLOADED L3L40 S L3 L5 0 S L3 0 S L3 FULL L6 STRUCTURE UPLOADED L7 L8 11 S L7 Ь9 801 S L7 FULL FILE 'HCAPLUS' ENTERED AT 22:13:42 ON 16 AUG 2006 L10 4 S L9 L11 2 S L10 AND GIBLIN, G?/AU => s l10 not l11 2 L10 NOT L11 L12 => s l12 and hall, a?/au 2142 HALL, A?/AU 0 L12 AND HALL, A?/AU L13 => s l12 and hurst, d?/au 286 HURST, D?/AU 0 L12 AND HURST, D?/AU L14 => s 112 and kilford, i?/an 0 KILFORD, I?/AN 0 L12 AND KILFORD, I?/AN L15 => s 112 and kilford, i?/au 10 KILFORD, I?/AU L16 0 L12 AND KILFORD, I?/AU => s 112 and lewell, x?/au 32 LEWELL, X?/AU 0 L12 AND LEWELL, X?/AU L17 => s 112 and taylor, a?/au 4316 TAYLOR, A?/AU 0 L12 AND TAYLOR, A?/AU L18 => s l12 and novelli, r?/au 38 NOVELLI, R?/AU 0 L12 AND NOVELLI, R?/AU L19 => d 112, ibib abs hitstr, 1-2 L12 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN 1971:76357 HCAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 74:76357 Noel bismethylene transfer to 2'-hydroxylated TITLE: isoflavones by dimethylsulfoxonium methylide: the reaction and its products Crombie, Leslie; Davies, John Salmon; Whiting, Donald AUTHOR(S): Dep. Chem., Univ. Coll. New South Wales, Cardiff, UK CORPORATE SOURCE: Journal of the Chemical Society [Section] C: Organic SOURCE: (1971), (2), 304-12 CODEN: JSOOAX; ISSN: 0022-4952 DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 74:76357

For diagram(s), see printed CA Issue.

Isoderritol isoflavone (I) reacted with excess ylide M/2S(O):CH2 to give a hydroxycyclopentene (II) and by-product decarboxyisor ϕ teno-nonic acid (III). The mechanism [ring cleavage, methylene transfer, and recyclization via the vinylcoumaranone (IV)] was disqussed. Acid rearrangement of II gave the stilbenoid cyclopentenone (V), and the dihydro derivative (VI) of II gave a hexacyclic compound (VII). Derritol isoflavone reacted similarly with Me2S(O):CH2. Equ‡molar amts. of I and Me2S(O):CH2 gave IV.

29517-95-1P 29558-29-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

29517-95-1 HCAPLUS RN

2-Cyclopenten-1-one, 3-(2-hydroxy-4,5-dimethoxyph@nyl)-2-(4-hydroxy-2-CN isopropyl-5-benzofuranyl)-, 3-acetate (8CI) (CA /INDEX NAME)

29558-29-0 HCAPLUS RN

2-Cyclopenten-1-one, 3-(2-hydroxy-4,5-dime/thoxyphenyl)-2-(4-hydroxy-2-CN isopropyl-5-benzofuranyl)-, diacetate (8C1) (CA INDEX NAME)

HCAPLUS COPYRIGHT 2006 ACS on STN L12 ANSWER 2 OF 2

ACCESSION NUMBER:

1970:435251 HQAPLUS

DOCUMENT NUMBER:

73:35251

TITLE:

Bis-methylene transfer to 2'-hydroxyisoflavones by dimethylsulfoxonium methylide

AUTHOR (S):

Crombie, Leslie; Davies, John Salmon; Whiting, D. A.

CORPORATE SOURCE:

Dep. Chem., Univ. Coll. Cardiff, Cardiff, UK Journal of the Chemical Society [Section] D: Chemical

SOURCE:

Communications (1970), 9, 535-6

CODEN: CCJDAO; ISSN: 0577-6171

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 73:35251

GI For diagram(s), see printed CA Issue.

AB Isoderritol isoflavone (I) with 1 and 5 moles Me2S+O-CH2 gave II and double methylene transfer product III, resp., and a small amount of IV. III was also obtained from II under similar reaction conditions, suggesting II as probable intermediate, and III rearranged readily in acid to V.

as probable intermediate, and III rearranged readily in acid to V. Dihydro derivative (VI) of III formed a monoacetate (VII) whose OH function was readily replaced in acids to give VIII or IX (R = H or D) depending on reaction conditions.

IT 29517-95-1P 29558-29-0P

RN 29517-95-1 HCAPLUS

CN 2-Cyclopenten-1-one, 3-(2-hydroxy-4,5-dimethoxyphenyl)-2-(4-hydroxy-2-isopropyl-5-benzofuranyl)-, 3-acetate (8CI) (CA INDEX NAME)

RN 29558-29-0 HCAPLUS

CN 2-Cyclopenten-1-one, 3-(2-hydroxy-4,5-dimethoxyphenyl)-2-(4-hydroxy-2-isopropyl-5-benzofuranyl)-, diacetate (8CI) (CA INDEX NAME)

=> d his

L1

(FILE 'HOME' ENTERED AT 22:05:35 ON 16 AUG 2006)

FILE 'REGISTRY' ENTERED AT 22:05:41 ON 16 AUG 2006 STRUCTURE UPLOADED

10508761

L2 0	S L1		
L3	STRUCTURE UPLOADED		
L4 0	S L3		
L5 0	S L3		
L6 0	S L3 FULL		
L7	STRUCTURE UPLOADED		
L8 11	S L7		
L9 801	S L7 FULL		
בדו.ם יטראסו	LUS' ENTERED AT 22:13:42 ON 1	16 NIG 2006	
-	S L9	10 A0G 2000	
	S L10 AND GIBLIN, G?/AU		
	S L10 NOT L11		
	S L12 AND HALL, A?/AU		
	S L12 AND HURST, D?/AU		
	S L12 AND KILFORD, I?/AN		
	S L12 AND KILFORD, I?/AU		
	S L12 AND LEWELL, X?/AU		
	S L12 AND TAYLOR, A?/AU		
	S L12 AND NOVELLI, R?/AU		
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COST IN U.S. DO	LLARS	SINCE FILE	
			SESSION
FULL ESTIMATED	COST	30.56	369.49
DISCOUNT AMOUNTS	S (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	

-3.00

-3.00

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FILE COVERS 1907-1966 FILE LAST UPDATED: 01 May 1997 (19970501/UP)

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